



The mitochondrial genome and phylogenetic characteristics of the Thick-billed Green-Pigeon, *Treron curvirostra*: the first sequence for the genus

Nan Xu^{1*}, Jiayu Ding^{1*}, Ziting Que¹, Wei Xu¹, Wentao Ye¹, Hongyi Liu¹

I College of Biology and the Environment, Nanjing Forestry University, Nanjing 210037, China

Corresponding author: Hongyi Liu (hongyi_liu@njfu.edu.cn)

Academic editor: G. Sangster | Received 30 October 2020 | Accepted 17 February 2021 | Published 2 June 2021

http://zoobank.org/E1EEC61A-4A09-41E2-A6A6-2B66F0D0B868

Citation: Xu N, Ding J, Que Z, Xu W, Ye W, Liu H (2021) The mitochondrial genome and phylogenetic characteristics of the Thick-billed Green-Pigeon, *Treron curvirostra*: the first sequence for the genus. ZooKeys 1041: 167–182. https://doi.org/10.3897/zookeys.1041.60150

Abstract

Members of the genus *Treron* (Columbidae) are widely distributed in southern Asia and the Indo-Malayan Region but their relationships are poorly understood. Better knowledge of the systematic status of this genus may help studies of historical biogeography and taxonomy. The complete mitochondrial genome of *T. curvirostra* was characterized, a first for the genus. It is 17,414 base pairs in length, containing two rR-NAs, 22 tRNAs, 13 protein coding genes (PCGs), and one D-loop with a primary structure that is similar to that found in most members of Columbidae. Most PCGs start with the common ATG codon but are terminated by different codons. The highest value of the Ka/Ks ratio within 13 PCGs was found in ATP8 with 0.1937, suggesting that PCGs of the mitochondrial genome tend to be conservative in Columbidae. Moreover, the phylogenetic relationships within Columbidae, which was based on sequences of 13 PCGs, showed that (*T. curvirostra* + *Hemiphaga novaeseelandiae*) were clustered in one clade, suggesting a potentially close relationship between *Treron* and *Hemiphaga*. However, the monophyly of the subfamilies of Columbidae recognized by the Interagency Taxonomic Information System could not be corroborated. Hence, the position of the genus *Treron* in the classification of Columbidae may have to be revised.

Keywords

Columbidae, genome sequencing, Ka/Ks ratio, mitochondrial DNA, phylogenetic tree

^{*} Authors contributed equally to this work.

Introduction

Mitochondrial DNA sequences can be reliable markers for studying the origin and phylogenetic relationships of species owing to its fast evolution rate, simple structure, light molecular weight, and maternal inheritance (Nabholz et al. 2016; Martins et al. 2019). Mitochondrial genomes of birds have a closed loop structure with lengths of 15,500–23,000 base pairs (bp) (Sammler et al. 2011; Xu et al. 2019; Wang et al. 2020). They typically contain 13 protein coding genes (PCGs), 22 transfer RNA genes (tRNAs), two ribosomal RNA genes (rRNAs), and one D-loop (Bensch and Härlid 2000; Sun et al. 2020), while some species were found to have duplicate regions (Eberhard and Wright 2016).

The pigeons and doves (family Columbidae) are widely distributed on all continents except Antarctica, ranging from tropical to temperate regions (Gibbs et al. 2001). The number of subfamilies of Columbidae differs among taxonomic authorities. Dickinson and Remsen (2013) recognize three subfamilies (Columbinae, Peristerinae, and Raphinae), whereas the Interagency Taxonomic Information System (ITIS) recognizes five subfamilies (Columbinae, Didunculinae, Gourinae, Otidiphabinae, and Treroninae), as well as 49 genera and more than 300 extant species (Integrated Taxonomic Information System 2020).

All species of green-pigeons (*Treron*) are listed as second-class national protected animals under China's Catalog of Wildlife of the Key State Protection. Most species in the genus are declining (Birdlife International 2018); however, only a few genetic resources are available for the genus *Treron* (e.g., Sorenson et al. 2003; Pereira et al. 2007; Hackett et al. 2008; Price et al. 2014; Claramunt and Cracraft 2015).

The Thick-billed Green-Pigeon *Treron curvirostra* (Gmelin, 1789) is mainly distributed in virgin, evergreen, broad-leaved, and secondary forests of the tropical and subtropical hilly zone in Southeast Asia and South Asia (Gibbs et al. 2001). Like most species of Columbidae, *T. curvirostra* feeds on seeds and fruits (Korzun et al. 2008). Members of this species have a medium-sized body and a colorful plumage (Korzun et al. 2008) distinguished by their grey head and green neck. The lower body is yellowish green, while the wing is nearly black, with a yellow feather margin and a distinct yellow wing spot. The central tail feathers are green, while the remaining feathers are gray with black secondary end spots (Korzun et al. 2008; Nair 2010). At present, only few studies have focused on *T. curvirostra*: Nair (2010) discussed the zoogeography.

To understand the systematic position of the genus *Treron* among Columbidae, we sequenced and characterized the first complete mitochondrial genome sequence of *T. curvirostra*. We compared the complete mitochondrial genome of *T. curvirostra* with that of 33 other pigeons and doves and determined its genetic structural characteristics. In addition, we used 13 protein-coding genes (PCGs) to reconstruct a phylogenetic tree, which we use to infer the taxonomic position of the species and illuminate the phylogenetic relationships among species of Columbidae.

Materials and methods

Sample collection and DNA extraction

This study was authorized by Nanjing Forestry University. The youngest tail feathers of a male Thick-billed Green-Pigeon *T. curvirostra* were collected from an individual rescued from a net that was used to prevent birds from stealing fruit at the Xieyang peak of Dali City, Yunnan Province, China. The bird was identified as *T. curvirostra* based on its morphological characters (Gibbs et al. 2001). After sample collection, the bird was released. The tail feather samples were transported to the Laboratory of Animal Molecular Evolution at the Nanjing Forestry University and stored at -80 °C. The tubules were cut and the pulp was removed for genomic DNA extraction using the FastPure Cell/Tissue Isolation Mini kit (Vazyme Biotechnology Co., Ltd., Nanjing, China) and stored at -20 °C for later use.

PCR amplification and sequencing

Primers were designed based on the mitochondrial gene sequences of *Streptopelia decaocto*, *Hemiphaga novaeseelandiae*, and *Columba hodgsonii* (GenBank accession numbers KY827036, EU725864, and MN919176,respectively) using DNASTAR software (DNASTAR, USA; Burland 2000). Primer sequences are listed in Table 1. The PCR reaction volume was 25 μ L, which included 1 μ L of template DNA, 12.5 μ L of the 2×Rapid Taq Master Mix (Vazyme Biotech Co., Ltd, Nanjing, China), 1 μ L per primer, and 9.5 μ L double-distilled (dd)H₂O. The PCR reaction procedure consisted of a pre-denaturation at 95 °C for 3 min, a denaturation at 95 °C for 15 s, an annealing at 50 °C to 60 °C for 15 s, which was adjusted according to the primers' own conditions, an extension at 72 °C for 2 min, cycling 35 times, and a final extension at 72 °C for

Table 1. Primers used for amplification of the *T. curvirostra* mitogenome.

Fragment	Region	Primer pair	Primer sequence (5' - 3')
DG 1	COI-COII	DG 1F	CACTCAGCCATCTTACCT
		DG 1R	ACAGATTTCTGAGCATTGGC
DG 2	COII-ND4	DG 2F	CCAATCCGCATCATCGTC
		DG 2R	GGTTTCCTCATCGTGTGA
DG 3	ND4-ND5	DG 3F	CAGCCTCCTAATTGCCAC
		DG 3R	GTAGGGCGGAGACTGGAG
DG 4	ND5-Cyt b	DG 4F	ACAGGGCCGAGCAGAAGC
		DG 4R	TAGGAAGTATCACTCTGG
DG 5	Cyt b-12S rRNA	DG 5F	GCAGGCCTCACCATTATCC
		DG 5R	GTTAATTACTGCTGAGTACC
DG 6	12S rRNA-16S rRNA	DG 6F	GCTGGCATCAGGCACGCC
		DG 6R	TGGGTCTGGTTACTGTTA
DG7	16S rRNA-ND2	DG 7F	CGGTTGGGGCGACCTTGGAG
		DG 7R	AGAGTGGGAGGAGTAGGGC
DG 8	ND2-COI	DG 8F	AGCAGCCACAATCATGGC
		DG 8R	ATAGATTTGGTCATCTCC

5 min. The PCR products were detected by a 1% agarose gel electrophoresis, and then sent to Tsingke Biotech Co., Ltd. (Nanjing, China), where the original primers were used for the bidirectional sequencing.

Sequence analysis

By comparing and identifying the DNA sequence of each mitochondrial gene in other pigeon families, the range and location of *T. curvirostra*'s mitochondrial genes were annotated. Hence, the complete mitochondrial genome sequence was used to predict the transcriptional direction of each gene component using the Improved de novo Metazoan Mitochondrial Genome Annotation (MITOS) platform (Bernt et al. 2013). The annotated mitochondrial genome sequence of *T. curvirostra* was submitted to GenBank (accession number MT535857). The mitochondrial ring structure was plotted, and 22 tRNA clover two-dimensional structures were predicted using programs, such as the comparative genomics (CG) View Server and the tRNAscan-Se (Stothard and Wishart 2005; Lowe and Chan 2016). Composition skew was calculated according to the following formulae: AT-skew = (A-T)/(A+T) and GC-skew = (G-C)/(G+C) (Perna and Kocher 1995). Moreover, the relative synonymous codon usage (RSCU) frequency and the ratio of the number of nonsynonymous substitutions per nonsynonymous site to the number of synonymous substitutions per synonymous site (Ka/Ks) of 13 PCGs of Columbidae were calculated using MEGA7 (Kumar et al. 2016), while the RSCU comparison graph was drawn by PhyloSuite (Zhang et al. 2020).

Phylogenetic analysis

We used a concatenated set of base sequences of the 13 PCGs from 34 pigeons and doves to investigate the phylogenetic position of T. curvirostra (Table 2). Yellowthroated Sandgrouse (Pterocles gutturalis Smith, 1836) was used as an outgroup. All operations were performed in the PhyloSuite software package (Zhang et al. 2020). The sequences were aligned in batches using MAFFT software (Katoh et al. 2002). ModelFinder was used to partition the codons and identify the best substitution model for the phylogenetic analyses (Kalyaanamoorthy et al. 2017). Phylogenetic trees were constructed with Bayesian inference (BI) and maximum-likelihood (ML) (Yang 1994; Huelsenbeck and Ronquist 2001). The best substitution model of BI was selected according to codon 1, 2 and 3, while the model of ML was determined by the automatic partitioning (Table 3). For the BI tree, Markov chains were run for one million generations and were sampled every 100 generations. The majority-rule consensus trees were estimated by combining the results from duplicated analyses, while discarding the first 25% of generations. Besides, we checked for nuclear copies of mitochondrial sequences (numts) and possible chimerism (Sangster et al. 2016; Sangster and Luksenburg 2020).

Table 2. Summary of the mitogenomes used in the analyses.

Family	Subfamily	Genus	Species	Accession
Columbidae	Columbinae	Geopelia	Geopelia cuneata	MN930521.1
			Geopelia striata	MG590276.1
		Trugon	Trugon terrestris	MG590263.1
		Caloenas	Caloenas maculata	KX902249.1
			Caloenas nicobarica	MG590264.1
		Streptopelia	Streptopelia tranquebarica	MT535858
			Streptopelia orientalis	KY827037.1
			Streptopelia decaocto	KY827036.1
			Streptopelia chinensis	KP273832.1
		Columba	Columba hodgsonii	MN919176.1
			Columba janthina	LC541479.1
			Columba jouyi	KX902247.1
			Columba livia	KP319029.1
			Columba rupestris	KX902246.1
		Ectopistes	Ectopistes migratorius	KC489473.1
		Patagioenas	Patagioenas fasciata	KX902240.1
		Leptotila	Leptotila verreauxi	HM640214.1
		Zenaida	Zenaida macroura	KX902235.1
			Zenaida auriculata	HM640211.1
		Geotrygon	Geotrygon violacea	HM640213.1
		Turtur	Turtur tympanistria	HM746793.1
		Columbina	Columbina picui	MN356335.1
		Chalcophaps	Chalcophaps indica	HM746789.1
	Treroninae	Alopecoenas	Alopecoenas salamonis	KX902250.1
		Hemiphaga	Hemiphaga novaeseelandiae	EU725864.1
	Gourinae	Goura	Goura cristata	MG590273.1
			Goura sclaterii	MG590285.1
			Goura scheepmakeri	MG590282.1
			Goura victoria	MG590299.1
		Pezophaps	Pezophaps solitaria	KX902238.1
		Raphus	Raphus cucullatus	KX902236.1
	Otidiphabinae	Otidiphaps	Otidiphaps nobilis	MG590265.1
	Didunculinae	Didunculus	Didunculus strigirostris	MG590266.1
Pteroclidae		Pterocles	Pterocles gutturalis	MN356147.1

Results and discussion

Mitochondrial genome structure and organization

The mitochondrial genome of the Thick-billed Green-Pigeon was found to be 17,414 bp in length, which agrees with the length of most of the other sequenced species of pigeons and doves (Table 4, Table 5, Table 6) (Pereira et al. 2007; Zhang et al. 2015). In addition, the base composition of *T. curvirostra* was found to be A = 30.32%, G = 13.61%, T = 24.83%, and C = 31.24%), where the A+T content (55.15%) was higher than the G+C content (44.85%) and is similar to other birds in Columbidae (Table 5 and Table 6) (Huang et al. 2016; Jang et al. 2016). Moreover, the genome had a closed circular ring structure, containing 22 tRNAs, 2 rRNAs, 13 PCGs, and one D-loop. The ND6 gene and the other 8 tRNAs (tRNA-Gln, tRNA-Ala, tRNA-

Table 3. The best substitution models for Bayesian inference (BI) and maximum-likelihood (ML) analyses.

		ND1	ND2	COI	COII	ATP6	ATP8	COIII	ND3	ND4L	ND4	ND5	Cyt b	ND6
BI	Codon 1	SYM +	GTR+	SYM +	SYM +	GTR+	GTR	SYM +	SYM +	GTR+	GTR+	GTR +	SYM +	GTR
		I + G4	F + I +	I + G4	I + G4	F + I +	+ F +	I + G4	I + G4	F + I +	F + I +	F + I +	I + G4	+ F +
			G4			G4	G4			G4	G4	G4		G4
	Codon 2	GTR+	HKY +											
		F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +
		G4												
	Codon 3	GTR+	GTR+	GTR+	GTR +	GTR +	GTR+	GTR+	GTR+	GTR+	GTR+	GTR +	GTR+	GTR
		F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	F + I +	+ F +
		G4												
ML		TVM +	TVM +	TIM2	TIM2	TVM +	TN +	TIM2	TIM2	TIM2	TVM +	TVM +	TIM2	TIM2
		F + R4	F + R4	+ F + I	+ F + I	F + R4	F + I +	+ F + I	+ F + I	+ F + I	F + R4	F + R4	+ F + I	+ F + I
				+ G4	+ G4		G4	+ G4	+ G4	+ G4			+ G4	+ G4

Table 4. Mitochondrial genetic composition of *T. curvirostra*.

Gene	Strand	Position	Anticodon	Size (bp)	Start codon		Intergenic length
tRNA-Phe	Н	1–68	GAA	68			0
12S rRNA	Н	69-1041		973			0
tRNA-Val	Н	1042-1114	UAC	73			0
16S rRNA	Н	1115-2703		1589			0
tRNA-Leu	Н	2704-2777	UAA	74			12
ND1	Н	2790-3755		966	ATG	AGA	17
tRNA-Ile	H	3773-3844	GAU	72			5
tRNA-Gln	L	3850-3920	UUG	71			0
tRNA-Met	Н	3921-3988	CAU	68			0
ND2	Н	3989-5027		1039	ATG	T	0
tRNA-Trp	Н	5028-5098	UCA	71			1
tRNA-Ala	L	5100-5168	UGC	69			2
tRNA-Asn	L	5171-5243	GUU	73			2
tRNA-Cys	L	5246-5313	GCA	68			0
tRNA-Tyr	L	5314-5384	GUA	71			1
COI	Н	5386-6936		1551	ATG	AGG	0
tRNA-Ser	L	6937-7001	UGA	65			2
tRNA-Asp	H	7004-7072	GUC	69			1
COII	Н	7074–7757		684	ATG	TAA	1
tRNA-Lys	Н	7759–7828	UUU	70			1
ATP8	Н	7830-7997		168	ATG	TAA	0
ATP6	Н	7988-8671		684	ATG	TAA	-1
COIII	H	8671-9454		784	ATG	T	0
tRNA-Gly	Н	9455-9523	UCC	69			0
ND3	H	9524-9875		352	ATT	TAA	1
tRNA-Arg	H	9877-9945	UCG	69			1
ND4L	Н	9947-10243		297	ATG	TAA	-7
ND4	H	10237-11614		1378	ATG	T	0
tRNA-His	Н	11615-11683	GUG	69			0
tRNA-Ser	Н	11684-11749	GCU	66			0
tRNA-Leu	Н	11750-11819	UAG	70			0
ND5	Н	11820-13637		1818	ATG	AGA	8
Cyt b	Н	13646-14788		1143	ATG	TAA	0
tRNA-Thr	Н	14789–14856	UGU	68			6
tRNA-Pro	L	14863-14932	UGG	70			4
ND6	L	14937-15458		522	ATG	TAG	3
tRNA-Glu	L	15462-15532	UUC	71			0
D-loop		15553-17414		1862			0

Region	A%	T%	AT-skew	G%	С%	GC-skew
whole mitogenome	30.32	24.83	0.100	13.61	31.24	-0.393
PCGs	29.46	24.56	0.091	12.23	33.76	-0.468
rRNAs	32.75	21.19	0.214	19.05	27.01	-0.173
tRNAs	32.33	25.16	0.125	16.95	25.55	-0.203
D-loop	30.45	31.31	-0.014	11.92	26.32	-0.376

Table 5. Composition and skewness in mitochondrial genome of *T. curvirostra*.

Table 6. Nucleotide composition indices in different regions of mitogenomes of Columbidae.

Species	GenBank no	Whole		Protein coding	Ribosomal RNA			
		mitogenome		genes				
		Length (bp)	AT (%)	Length (bp)	AT (%)	Length (bp)	AT (%)	
Goura sclaterii	MG590285.1	18242	54.13	11386	52.99	2571	53.21	
Treron curvirostra	MT535857	17414	55.16	11386	54.02	2562	53.94	
Columba hodgsonii	MN919176.1	17477	54.55	11385	53.82	2557	53.23	
Trugon terrestris	MG590263.1	17405	55.62	11395	54.94	2569	55.86	
Didunculus strigirostris	MG590266.1	17389	54.94	11390	54.32	2569	55	
Geopelia striata	MG590276.1	17354	54.83	11383	53.78	2565	54.07	
Otidiphaps nobilis	MG590265.1	17346	55.83	11382	55.2	2581	54.75	
Hemiphaga novaeseelandiae	EU725864.1	17264	54.89	11386	54.1	2575	54.52	
Caloenas nicobarica	MG590264.1	17178	55.13	11386	54.83	2567	53.64	
Leptotila verreauxi	HM640214.1	17176	54.1	11383	53.34	2556	53.4	
Alopecoenas salamonis	KX902250.1	17141	55.18	11381	54.78	2569	54.81	
Streptopelia orientalis	KY827037.1	17102	53.41	11386	52.86	2561	53.58	
Raphus cucullatus	KX902236.1	17092	56.08	11385	55.87	2566	54.6	
Ectopistes migratorius	KC489473.1	17026	54.52	11383	54.3	2596	52.97	
Patagioenas fasciata	KX902239.1	16970	54.51	11384	54.02	2550	53.8	
Geotrygon violacea	HM640213.1	16864	55.04	11383	54.27	2561	53.85	
Zenaida auriculata	HM640211.1	16781	53.29	11380	52.71	2567	52.83	

Asn, tRNA-Cys, tRNA-Tyr, tRNA-Ser (UGA), tRNA-Pro, and tRNA-Glu) were transcribed from the light (L)-strand, while the other genes were transcribed from the heavy (H)-strand (Fig. 1, Table 4). In addition, two pairs of overlapping regions among the ATP6/COIII and ND4L/ND4 were found, with an overlapping region of ATP6/COIII being one bp and the overlapping region of ND4L/ND4 being seven bp. Furthermore, 18 intergenic spacers were observed between the mitochondrial regions with lengths between -7 and 17 bp. Among all these intergenic spacers, the shortest was -7 bp (found between ND4L and ND4), while the longest was 17 bp (found between ND1 and tRNA-Ile).

The PCGs

The total length of the PCGs was 11,386 bp, which is consistent with the average length of PCGs found in Columbidae (Table 5). The base composition of PCGs was A = 29.46%, G = 12.23%, T = 24.56%, and C = 33.76%, while the A+T content (54.01%) was slightly higher than the G+C content (45.99%). The AT-skew of *T. curvirostra* was positive, while the GC-skew was negative (Table 5). Furthermore, the PCG regions of *T. curvirostra* contained genes coding for cytochrome b (Cytb), two

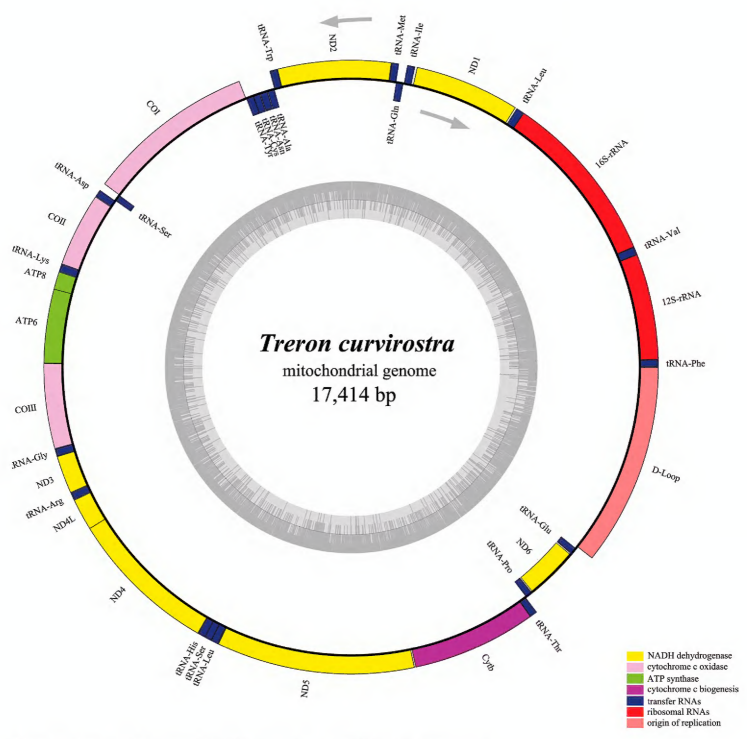


Figure 1. Circular map of the *T. curvirostra* mitochondrial genome.

ATPases (ATP6 and ATP8), three cytochrome c oxidases (COI, COII, and COIII), and seven NADH dehydrogenases (ND1-6 and ND4L). With the exception of ND3 (which had ATT as its start codon), all the other PCGs had ATG as a start codon. Six PCGs had the complete stop codon of TAA, while four PCGs had the other complete stop codons of AGA (ND1 and ND5), AGG (COI), and TAG (ND6). ND2, ND4, and COIII had the incomplete stop codon of T (Table 4). The RSCU of *T. curvirostra* is illustrated in Fig. 2, where Leu1 had the highest concentration and Cys had the lowest. In addition, Met only had AUG, while the other seven regions had four codons. With *T. curvirostra* as a baseline, the Ka/Ks ratio (Hurst 2002) of the 13 PCGs in 17 species of doves were all less than 1, with the highest Ka/Ks ratio (0.1937) in ATP8 and the lowest ratio (0.0243) in COI (Fig. 3). Hence, it seems that evolution tended to be conservative and maintained the generated protein (Hanada et al. 2007).

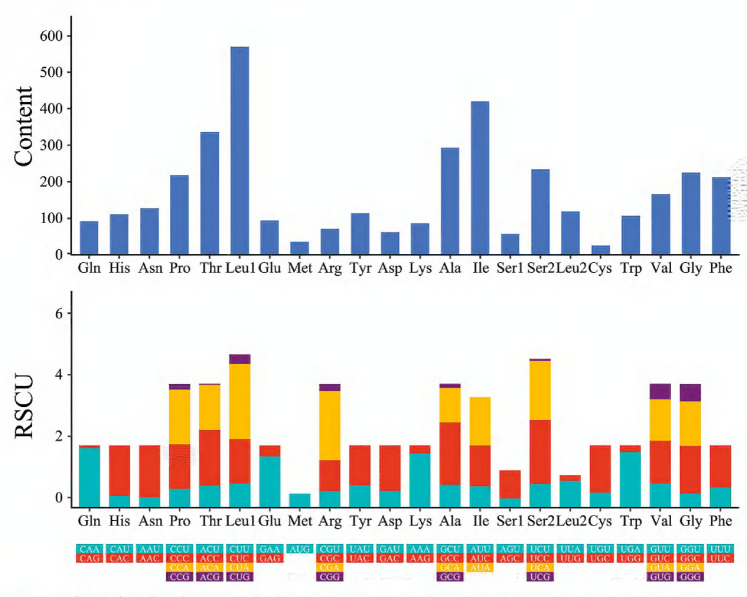


Figure 2. Codon distribution and relative synonymous codon usage in *T. curvirostra* mitogenome.

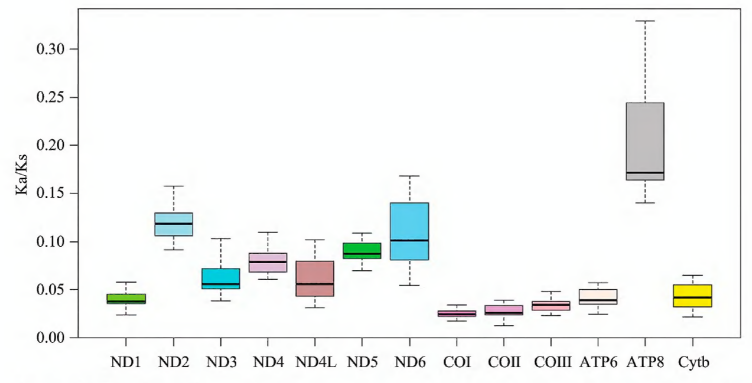


Figure 3. The ratio of the number of nonsynonymous substitutions per nonsynonymous site to the number of synonymous substitutions per synonymous site of 13 PCGs among 17 species of pigeons and doves. *T. curvirostra* was set as a baseline.

Transfer RNAs, ribosomal RNAs, and the D-loop

The mitogenome of *T. curvirostra* contained 22 tRNAs with lengths ranging from 65 bp (tRNA-Ser (UGA)) to 74 bp (tRNA-Leu (UAA)), which is similar to that in the mitogenomes of other pigeons and doves (Zhang et al. 2015). Moreover, the total length of the tRNAs was 1,534 bp, with an A+T content of 57.50%, a G+C content of 42.50%, an AT-skew of 0.1247, and a GC-skew of -0.2025 (Table 5). Among all the secondary structures of the 22 tRNA genes from the *T. curvirostra* mitochondrial genome, with the exception of tRNA-Ser (GCU), all had a typical cloverleaf structure (Fig. 4).

The total size of the two rRNAs was 2,562 bp, with an A+T content of 53.94%, an AT-skew of 0.2142, and a GC-skew of -0.1729 (Table 5). The 12S rRNA was 973 bp in length and was located between tRNA-Phe and tRNA-Val, while the 16S rRNA was 1,589 bp in length and was located between tRNA-Val and tRNA-Leu (UAA).

A D-loop was found between tRNA-Glu and tRNA-Phe, and was 1,862 bp in length, with a A+T content of 61.76%, an AT-skew of -0.0139, and a GC-skew of -0.3764 (Table 5). Duplication and rearrangement of the avian mitochondrial genomes is common, but *T. curvirostra* had only one D-loop, which is similar to that present in other known mitogenomes of Columbidae (Pacheco et al. 2011; Eberhard and Wright 2016; Bruxaux et al. 2018).

Phylogenetic analysis

Although the topology of ML tree and BI tree were similar to each other, they differed with respect to the phylogenetic position of T. curvirostra. Treron curvirostra clustered with Hemiphaga novaeseelandiae (Gmelin, 1789) in the BI tree, whereas it did not cluster with any species in the ML tree (Fig. 5). Therefore, we tested for the presence of the numts and chimerism. All these tests were negative, indicating the validity of *T. curvirostra* mitogenome. The phylogenetic trees also highlighted the stable relationships among the same genera within Columbidae, which was consistent with previous studies from analyses of mitochondrial and nuclear genes (Kan et al. 2010; Pacheco et al. 2011; Hung et al. 2013; Mlíkovský 2016; Soares et al. 2016; Kretschmer et al. 2020; Liu et al. 2020) (Fig. 5). However, the phylogenetic analysis did not support the arrangement of pigeons into five subfamilies (Columbinae, Didunculinae, Gourinae, Otidiphabinae, and Treroninae) as recognized by ITIS. Caloenas, Geopelia, and Trugon terrestris (which were placed in Columbinae by ITIS) clustered with species from other subfamilies in our phylogenies (Fig. 5). The most likely cause might be that the original classification system was based mainly on patterns of overall similarity in morphology which may not accurately reflect phylogenetic relationships. Similar contradictions between overall similarity and phylogeny have also been found in other groups of birds, including terns (Bridge et al. 2005), rails (Sangster et al. 2015), nightjars (Han et al. 2010), eagles (Lerner and Mindell 2005), laughing thrushes (Luo et al. 2008), and chats and flycatchers (Sangster et al. 2010). Our results indicate that

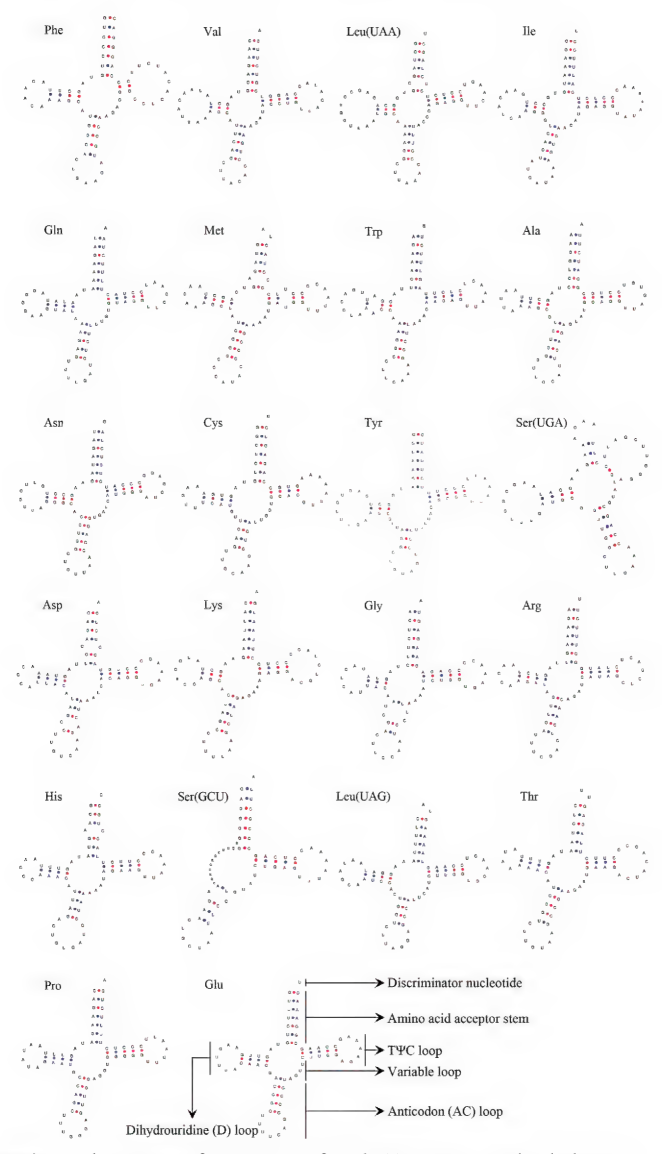


Figure 4. Secondary structure of 22 tRNA genes from the *T. curvirostra* mitochondrial genome.

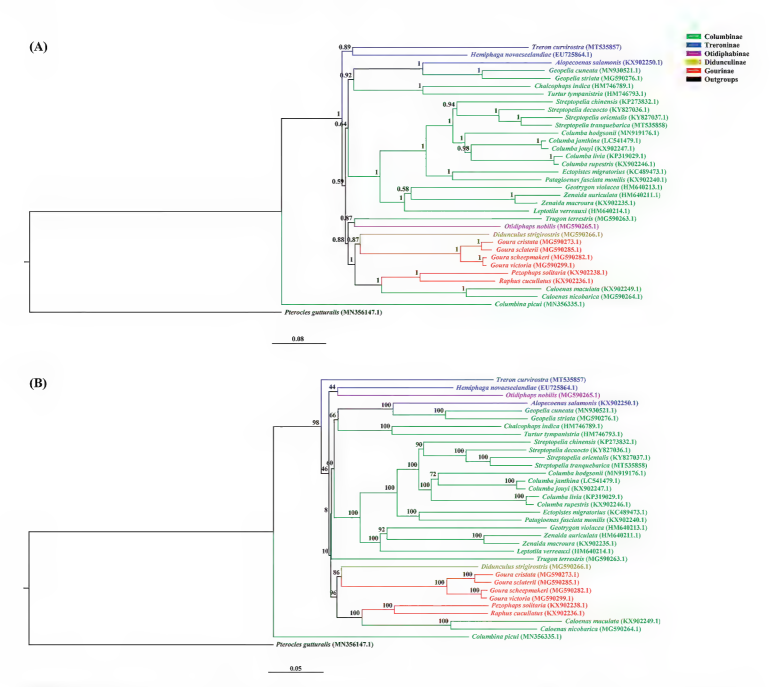


Figure 5. Mitogenomic phylogeny of 34 Columbidae species and an outgroup (*Pterocles gutturalis*) based on 13 PCGs using the Bayesian inference (**A**) and maximum likelihood (**B**) methods. Different colors indicated different subfamilies.

the subfamily classification of Columbidae may not accurately reflect historical relationships and may need to be revised. However, the poor branch support of basal clades of Columbidae precludes such a revision at present. Clearly, future attempts to resolve the phylogeny of Columbidae with confidence should include a suitable set of nuclear markers.

Acknowledgements

The authors declare no competing interest exists. This study was supported by the National Natural Science Foundation of China (No. 31800453), the Priority Academic Program Development of Jiangsu Higher Education Institutions (PAPD), and the Innovation and Entrepreneurship Training Program for College Students of China (202010298035Z).

References

- Bensch S, Härlid A (2000) Mitochondrial genomic rearrangements in songbirds. Molecular Biology and Evolution 17: 107–113. https://doi.org/10.1093/oxfordjournals.molbev.a026223
- Bernt M, Donath A, Jühling F, Externbrink F, Florentz C, Fritzsch G, Pütz J, Middendorf M, Stadler PF (2013) MITOS: improved de novo metazoan mitochondrial genome annotation. Molecular Phylogenetics and Evolution 69: 313–319. https://doi.org/10.1016/j. ympev.2012.08.023
- BirdLife International (2018) *Treron curvirostra*. The IUCN Red List of Threatened Species 2018: e.T22691160A130177198. https://doi.org/10.2305/IUCN.UK.2018-2.RLTS. T22691160A130177198.en [accessed 18 January 2021]
- Bridge ES, Jones AW, Baker AJ (2005) A phylogenetic framework for the terns (Sternini) inferred from mtDNA sequences: implications for taxonomy and plumage evolution. Molecular Phylogenetics and Evolution 35: 459–469. https://doi.org/10.1016/j.ympev.2004.12.010
- Bruxaux J, Gabrielli M, Ashari H, Prŷs-Jones R, Joseph L, Milá B, Besnard G, Thébaud C (2018) Recovering the evolutionary history of crowned pigeons (Columbidae: *Goura*): Implications for the biogeography and conservation of New Guinean lowland birds. Molecular Phylogenetics and Evolution 120: 248–258. https://doi.org/10.1016/j. ympev.2017.11.022
- Burland TG (2000) DNASTAR's Lasergene sequence analysis software. Methods in Molecular Biology 132: 71–91. https://doi.org/10.1385/1-59259-192-2:71
- Claramunt S, Cracraft J (2015) A new time tree reveals Earth history's imprint on the evolution of modern birds. Science Advances 1: e1501005. https://doi.org/10.1126/sciadv.1501005
- Eberhard JR, Wright TF (2016) Rearrangement and evolution of mitochondrial genomes in parrots. Molecular Phylogenetics and Evolution 94: 34–46. https://doi.org/10.1016/j. ympev.2015.08.011
- Gibbs D, Barnes E, Cox J (2001) Pigeons and doves: a guide to the pigeons and doves of the world. Pica Press, Robertsbridge.
- Hackett SJ, Kimball RT, Reddy S, Bowie RCK, Braun EL, Braun MJ, Chojnowski JL, Cox WA, Han K-L, Harshman J, Huddleston CJ, Marks BD, Miglia KJ, Moore WS, Sheldon FH, Steadman DW, Witt CC, Yuri T (2008). A phylogenomic study of birds reveals their evolutionary history. Science 320: 1763–1768. https://doi.org/10.1126/science.1157704
- Han K-L, Robbins MB, Braun MJ (2010) A multi-gene estimate of phylogeny in the nightjars and nighthawks (Caprimulgidae). Molecular Phylogenetics and Evolution 55: 443–453. https://doi.org/10.1016/j.ympev.2010.01.023
- Hanada K, Shiu SH, Li WH (2007) The nonsynonymous/synonymous substitution rate ratio versus the radical/conservative replacement rate ratio in the evolution of mammalian genes. Molecular Biology and Evolution 24: 2235–2241. https://doi.org/10.1093/molbev/msm152
- Huang ZH, Tu FY, Liu XH (2016) Determination of the complete mitogenome of Spotted Dove, *Spilopelia chinensis* (Columbiformes: Columbidae). Mitochondrial DNA Part A 27: 4224–4225. https://doi.org/10.3109/19401736.2015.1022750
- Huelsenbeck JP, Ronquist F (2001) MRBAYES: Bayesian inference of phylogenetic trees. Bioinformatics 17: 754–755. https://doi.org/10.1093/bioinformatics/17.8.754

- Hung CM, Lin RC, Chu JH, Yeh CF, Yao CJ, Li SH (2013) The de novo assembly of mitochondrial genomes of the extinct Passenger Pigeon (*Ectopistes migratorius*) with next generation sequencing. PLoS ONE 8: e56301. https://doi.org/10.1371/journal.pone.0056301
- Hurst LD (2002) The Ka/Ks ratio: diagnosing the form of sequence evolution. Trends in Genetics 18: 486–487. https://doi.org/10.1016/S0168-9525(02)02722-1
- Integrated Taxonomic Information System [ITIS] (2020) Columbidae. Taxonomic Serial No.: 177061. http://www.itis.gov
- Jang KH, Ryu SH, Kang SG, Hwang UW (2016) Complete mitochondrial genome of the Japanese Wood Pigeon, *Columba janthina* (Columbiformes, Columbidae). Mitochondrial DNA Part A 27: 2165–2166. https://doi.org/10.3109/19401736.2014.982608
- Kalyaanamoorthy S, Minh BQ, Wong TKF, von Haeseler A, Jermiin LS (2017) Modelfinder: fast model selection for accurate phylogenetic estimates. Nature Methods 14: 587–589. https://doi.org/10.1038/nmeth.4285
- Kan XZ, Li XF, Zhang LQ, Chen L, Qian CJ, Zhang XW, Wang L (2010) Characterization of the complete mitochondrial genome of the Rock Pigeon, *Columba livia* (Columbiformes: Columbidae). Genetics and Molecular Research 9: 1234–1249. https://doi.org/10.4238/vol9-2gmr853
- Katoh K, Misawa K, Kuma K, Miyata T (2002) MAFFT: a novel method for rapid multiple sequence alignment based on fast Fourier transform. Nucleic Acids Research 30: 3059–3066. https://doi.org/10.1093/nar/gkf436
- Korzun LP, Erard C, Gasc JP, Dzerzhinsky FJ (2008) Bill and hyoid apparatus of pigeons (Columbidae) and sandgrouse (Pteroclididae): a common adaptation to vegetarian feeding? Comptes Rendus Biologies 331: 64–87. https://doi.org/10.1016/j.crvi.2007.10.003
- Kretschmer R, Furo IO, Gomes AJB, Kiazim LG, Gunski RJ, Del Valle Garnero A, Pereira JC, Ferguson-Smith MA, Corrêa de Oliveira EH, Griffin DK, Freitas TRO, O'Connor RE (2020) A comprehensive cytogenetic analysis of several members of the family Columbidae (Aves, Columbiformes). Genes 11(6): e632. https://doi.org/10.3390/genes11060632
- Kumar S, Stecher G, Tamura K (2016) MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. Molecular Biology and Evolution 33: 1870–1874. https://doi.org/10.1093/molbev/msw054
- Lerner HRL, Mindell DP (2005) Phylogeny of eagles, Old World vultures, and other Accipitridae based on nuclear and mitochondrial DNA. Molecular Phylogenetics and Evolution 37: 327–346. https://doi.org/10.1016/j.ympev.2005.04.010
- Liu HY, Sun CH, Zhu Y, Zhang QZ (2020) Complete mitogenomic and phylogenetic characteristics of the Speckled Wood-pigeon (*Columba hodgsonii*). Molecular Biology Reports 47: 3567–3576. https://doi.org/10.1007/s11033-020-05448-w
- Lowe TM, Chan PP (2016) tRNAscan–SE on–line: search and contextual analysis of transfer RNA genes. Nucleic Acids Research 44: W54–W57. https://doi.org/10.1093/nar/gkw413
- Luo X, Qu YH, Han LX, Li SH, Lei FM (2008) A phylogenetic analysis of laughing thrushes (Timaliidae: *Garrulax*) and allies based on mitochondrial and nuclear DNA sequences. Zoologica Scripta 38: 9–22. https://doi.org/10.1111/j.1463-6409.2008.00355.x
- Martins G, Balbino E, Marques A, Almeida C (2019) Complete mitochondrial genomes of the *Spondias tuberosa* Arr. Cam and *Spondias mombin* L. reveal highly repetitive DNA sequences. Gene 720: 144026. https://doi.org/10.1016/j.gene.2019.144026

- Mlíkovský J (2016) The type species of the genus *Geotrygon* Gosse, 1847 (Aves: Columbidae). Zootaxa 4126: 138–140. https://doi.org/10.11646/zootaxa.4126.1.8
- Nabholz B, Lanfear R, Fuchs J (2016) Body mass-corrected molecular rate for bird mitochondrial DNA. Molecular Ecology 25: 4438–4449. https://doi.org/10.1111/mec.13780
- Nair MV (2010) Thick-billed Green-Pigeon *Treron curvirostra* in Similipal Hills, Orissa: an addition to the avifauna of peninsular India. Indian Birds 6: 19–20. http://www.indianbirds.in/pdfs/Nair_ThickbilledGreenPigeon.pdf
- Pacheco MA, Battistuzzi FU, Lentino M, Aguilar RF, Kumar S, Escalante AA (2011) Evolution of modern birds revealed by mitogenomics: timing the radiation and origin of major orders. Molecular Biology and Evolution 28: 1927–1942. https://doi.org/10.1093/molbev/msr014
- Pereira SL, Johnson KP, Clayton DH, Baker AJ (2007) Mitochondrial and nuclear DNA sequences support a Cretaceous origin of Columbiformes and a dispersal-driven radiation in the Paleogene. Systematic Biology 56: 656–672. https://doi.org/10.1080/10635150701549672
- Perna NT, Kocher TD (1995) Patterns of nucleotide composition at fourfold degenerate sites of animal mitochondrial genomes. Journal of Molecular Evolution 41: 353–358. https://doi.org/10.1007/BF01215182
- Reddy S, Kimball RT, Pandey A, Hosner PA, Braun MJ, Hackett SJ, Han KL, Harshman J, Huddleston CJ, Kingston S, Marks BD, Miglia KJ, Moore WS, Sheldon FH, Witt CC, Yuri T, Braun EL (2017) Why do phylogenomic data sets yield conflicting trees? Data type influences the avian tree of life more than taxon sampling. Systematic Biology 66: 857–879. https://doi.org/10.1093/sysbio/syx041
- Sammler S, Bleidorn C, Tiedemann R (2011) Full mitochondrial genome sequences of two endemic Philippine hornbill species (Aves: Bucerotidae) provide evidence for pervasive mitochondrial DNA recombination. BMC Genomics 12: e35. https://doi.org/10.1186/1471-2164-12-35
- Sangster G, Alström P, Forsmark E, Olsson U (2010) Multi-locus phylogenetic analysis of Old World chats and flycatchers reveals extensive paraphyly at family, subfamily and genus level (Aves: Muscicapidae). Molecular Phylogenetics and Evolution 57: 380–392. https://doi.org/10.1016/j.ympev.2010.07.008
- Sangster G, García-R JC, Trewick SA (2015) A new genus for the Lesser Moorhen *Gallinula angulata* Sundevall, 1850 (Aves, Rallidae). European Journal of Taxonomy 153: 1–8. https://doi.org/10.5852/ejt.2015.153
- Sangster G, Luksenburg JA (2020) Chimeric mitochondrial genomes: a hazard for phylogenetics and environmental DNA identification of fishes. Authorea (Online). https://doi.org/10.22541/au.160205226.65255244/v1
- Sangster G, Roselaar CS, Irestedt M, Ericson PGP (2016) Sillem's Mountain Finch *Leucosticte sillemi* is a valid species of rosefinch (*Carpodacus*, Fringillidae). Ibis 158: 184–189. https://doi.org/10.1111/ibi.12323
- Soares AER, Novak BJ, Haile J, Heupink TH, Fjeldså J, Gilbert MTP, Poinar H, Church GM, Shapiro B (2016) Complete mitochondrial genomes of living and extinct pigeons revise the timing of the columbiform radiation. BMC Evolutionary Biology 16: e230. https://doi.org/10.1186/s12862-016-0800-3
- Sorenson MD, Oneal E, García-Moreno J, Mindell DP (2003) More taxa, more characters: the Hoatzin problem is still unresolved. Molecular Biology and Evolution 20: 1484–1498. https://doi.org/10.1093/molbev/msg157

- Stothard P, Wishart DS (2005) Circular genome visualization and exploration using CGView. Bioinformatics 21: 537–539. https://doi.org/10.1093/bioinformatics/bti054
- Sun CH, Liu HY, Lu CH (2020) Five new mitogenomes of *Phylloscopus* (Passeriformes, Phylloscopidae): Sequence, structure, and phylogenetic analyses. International Journal of Biological Macromolecules 146: 638–647. https://doi.org/10.1016/j.ijbiomac.2019.12.253
- Wang E, Zhang D, Braun MS, Hotz-Wagenblatt A, Pärt T, Arlt D, Schmaljohann H, Bairlein F, Lei F, Wink M (2020) Can mitogenomes of the Northern Wheatear (*Oenanthe oenanthe*) reconstruct its phylogeography and reveal the origin of migrant birds? Scientific Reports 10: e9290. https://doi.org/10.1038/s41598-020-66287-0
- Xu N, Zhang QZ, Chen R, Liu HY (2019) The complete mitogenome of Red-collared Lorikeet (*Trichoglossus rubritorquis*) and its phylogenetic analysis. Mitochondrial DNA Part B 4: 3116–3117. https://doi.org/10.1080/23802359.2019.1667917
- Yang Z (1994) Maximum likelihood phylogenetic estimation from DNA sequences with variable rates over sites: approximate methods. Journal of Molecular Evolution 39: 306–314. https://doi.org/10.1007/BF00160154
- Zhang D, Gao FL, Jakovlić I, Zou H, Zhang J, Li WX, Wang GT (2020) PhyloSuite: An integrated and scalable desktop platform for streamlined molecular sequence data management and evolutionary phylogenetics studies. Molecular Ecology Resources 20: 348–355. https://doi.org/10.1111/1755-0998.13096
- Zhang RH, Xu MJ, Wang CL, Xu T, Wei D, Liu BJ, Wang GH (2015) The complete mitochondrial genome of the Fancy Pigeon, *Columba livia* (Columbiformes: Columbidae). Mitochondrial DNA 26: 162–163. https://doi.org/10.3109/19401736.2014.1003851